## **PCT**

## WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



#### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:

C07C 17/20, 19/08, B01J 3/04, 19/02

A1

(11) International Publication Number: WO 99/00344

(43) International Publication Date: 7 January 1999 (07.01.99)

US

(21) International Application Number: PCT/US98/12951

(22) International Filing Date: 22 June 1998 (22.06.98)

(30) Priority Data: 08/882,805 26 June 1997 (26.06.97)

(71) Applicant: ALLIEDSIGNAL INC. [US/US]; 101 Columbia Road, P.O. Box 2245, Morristown, NJ 07962-2245 (US).

(72) Inventors: TUNG, Hsueh, Sung; 16 Vassar Drive, Getzville, NY 14968 (US). MERKEL, Daniel, Christopher; 64 Glenmar Drive, West Seneca, NY 14224 (US). SMITH, Addison, Miles; 80 Berryman Drive, Amherst, NY 14226 (US).

(74) Agent: CRISS, Roger, H.; AlliedSignal Inc., (Law Dept., Attn: R. Fels), 101 Columbia Road, P.O. Box 2245, Morristown, NJ 07962-2245 (US).

(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

#### **Published**

With international search report.

(54) Title: PROCESS FOR PREPARING HYDROFLUOROCARBONS

#### (57) Abstract

A process for producing a hydrofluorocarbon (HFC) comprising: (a) providing a fluoropolymer-lined reactor, (b) adding a chlorinated organic compound in liquid phase and a fluorination agent to the reactor, and (c) reacting at least a portion of the chlorinated organic compound with at least a portion of the fluorination agent to produce the HFC.

#### FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
ВJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazi!	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Vict Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
СН	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

# PROCESS FOR PREPARING HYDROFLUOROCARBONS FIELD OF INVENTION

The present invention relates generally to the preparation of hydrofluorocarbons (HFCs). More specifically, this invention relates to a fluorination process that minimizes reactor corrosion and improves the yield and purity of the HFCs produced.

5

10

15

20

25

#### BACKGROUND OF THE INVENTION

Because hydrofluorocarbons (HFCs) do not deplete the ozone layer, they are becoming popular substitutes for chlorofluorocarbons (CFCs) and hydrochlorofluorocarbons (HCFCs) for use as heat transfer agents, blowing agents, and propellants. HFCs are typically prepared by fluorinating a chlorinated organic compound with a fluorination agent such as hydrogen fluoride in the presence of a fluorination catalyst. This reaction may be conducted in either the liquid or gas phase. Generally, the liquid phase fluorination is preferred because the reaction is controlled at relatively lower temperatures which results in less byproduct formation due to decomposition.

Liquid phase fluorination, however, uses and generates corrosive compounds, such as, for example, hydrogen fluoride, hydrogen chloride, and catalysts, which form superacids. These superacids tend to corrode the reactor in which the reaction is conducted, even reactors comprised of corrosion-resistant materials such as Inconel 600, NAR25-50MII, Hastelloy C, Hastelloy G-30, duplex stainless steel, and Hastelloy C-22. Corrosion of the reactor compromises the structural integrity of the reactor and reduces its useful life. Therefore, a need to minimize reactor corrosion exists.

One method of reducing such corrosion is taught in Japanese Kokai Patent Application Publication No. 233102(1995). In this publication, a method is

disclosed for the liquid phase fluorination of a chlorinated organic compound in a reactor made or lined with a fluorine resin. The method involves gaseous feeds of hydrogen fluoride and chlorinated organic compound. Because the process is restricted to gaseous feed streams, it is limited in the type of HFCs it can produce. Chlorinated organic compounds having two or more carbon atoms tend to decompose before reaching their gaseous state. For example, pentachloropropane tends to decompose significantly at temperature greater than 90°C while its boiling point is about 190°C. Thus, as a practical matter, the process disclosed in this publication can only be used to produce fluorinated methanes.

10

5

The aforementioned Japanese publication also states that when heat transfer through the reactor is necessary, which is usually the case in liquid phase fluorination, the fluorine resin liner should be applied using a molding method. The only molding method identified therein is rotary-baked molding.

15

Generally, reactors having a molded liner, such as a rotary-baked or sprayed-on liner, are not suitable for large-scale commercial production. Reactors having such liners must be baked in large kilns or ovens, which are expensive and frequently unavailable. Indeed, fitting a large reactor, for example, greater than about a 1,000 gallons, with a baked liner is impractical.

20

A molded liner not only imposes practical limitations on the reactor, but also introduces structural limitations. It has been found that molded liners tend to be permeable and, under high pressures and over time, reactants tend to penetrate the liner and develop pressure between the liner and the reactor wall. This pressure causes the liner to blister, and eventually the liner comes loose. The problem of liner penetration is exacerbated by the absence of weep holes in a molded-liner reactor. Ordinarily, weep holes allow reactants that penetrate the liner to escape from the reactor. A molded liner, however, generally cannot be used in a reactor with weep holes. When applying a molded liner, a fluid fluoropolymer is applied to the reactor wall, and, thus, holes in the reactor wall will result in holes in the molded liner. Holes in the liner obviously compromise the reactor's ability to be pressurized. Therefore, while a rotary-baked, fluorine-

30

resin liner may minimize reactor corrosion, its structural limitations nevertheless limit the reactor's useful life.

Therefore, a need exists for a commercially viable method of producing a wide range of HFCs while minimizing reactor corrosion. The present invention fulfills this need among others.

## DESCRIPTION OF THE INVENTION AND PREFERRED EMBODIMENTS

The present invention is directed to a fluorination process which uses a commercially viable and durable fluoropolymer-lined reactor, as described hereafter, to produce a wide variety of hydrofluorocarbons (HFCs), including those having two or more carbon atoms.

In one embodiment, the process comprises: (a) providing reactor means for effecting fluorination in which at least a portion of the reactor means is fluoropolymer-lined; (b) adding a fluorination agent and a liquid chlorinated organic compound to the reactor means; and (c) reacting in the liquid phase at least a portion of the starting materials to produce a HFC. Since this embodiment adds the chlorinated organic compound to the reactor in *liquid* phase rather than gas phase, compounds having two or more carbon atoms can be used without decomposing.

20

5

10

15

Another embodiment of the process comprises: (a) providing a reactor with a loose, fluoropolymer liner; and (b) reacting at least a portion of a chlorinated organic compound with at least a portion of a fluorination agent in the reactor to produce an HFC. A loose, fluoropolymer liner, as described hereafter, provides a reactor with a chemically-resistant barrier that is more durable then molded-type, fluoropolymer liners. More specifically, a loose liner tends to be less porous than a molded liner and, thus, tends to maintain a seal for a longer time.

25

Yet another embodiment of the process comprises: (a) providing a fluoropolymer-lined reactor having weep holes; and (b) reacting at least a portion of a chlorinated organic compound with at least a portion of a fluorination agent in

the reactor to produce an HFC. The weep-holes allow those reactants that permeate the fluoropolymer-lined reactor to be vented or otherwise removed. This prevents reactants from building up pressure between the liner and the reactor wall and forming blisters in the liner.

In a preferred embodiment, the process comprises a combination of the above embodiments.

5

10

15

20

25

30

An important aspect of the present invention is the use of a fluoropolymer liner on at least a portion of the reactor means. As used herein, the terms "fluorinated polymer" and "fluoropolymer" are used synonymously and broadly refer to any polymer, copolymer or blend of polymers having a fluoride atom in at least one of the monomers. Preferred materials include, for example, polytetrafluoroethylene, poly(vinylidene fluoride), ethylene-tetrafluoroethylene polymer, ethylene-hexafluoropropylene polymer, tetrafluoroethylene-hexafluoropropylene polymer, tetrafluoroethylene-hexafluoropropylene polymer, any modified version of the above-mentioned polymers, and blends of two or more thereof. The polytetrafluoroethylene liner or its modified version is more preferred.

The reactor means for facilitating fluorination may comprise any apparatus conventionally used for preparing HFCs by liquid phase fluorination. An example of a satisfactory apparatus for this purpose is one consisting of a single reaction vessel, such as an autoclave, to which the starting materials can be added, in liquid or gaseous form, and heated or cooled well enough to keep the reaction temperature at a set temperature. The reaction vessel should promote contact among the reactants by a suitable mixing method and be capable of sustaining reaction pressures up to about 300 psi or whatever the maximum reaction pressure is expected to be. Because the reaction typically takes place under pressure, the reactor vessel is generally comprised of metal or other structurally rigid material. Suitable materials include, for example, carbon steel, stainless steel, Inconel alloy, Monel alloy, Hastelloy, or other type of a structurally suitable alloy.

In the preferred embodiment, the reactor wall has at least one weep hole, and preferably a plurality of weep holes. Weep holes allow reactants that

permeate the liner to be vented or otherwise removed from between the liner and the reactor wall. The number and size of the weep holes may vary according to reactor size and other variables, although a weep hole having a diameter from about 1/8 to about 1/2 inch is generally preferred. A reactor having one or more weep holes requires that the liner have adequate strength to span the weep hole under reaction pressures.

5

10

15

20

25

**30** 

The reaction means may comprise also a catalyst stripper which is operatively connected to the reaction vessel. A catalyst stripper typically consists of a condenser and a packed column, and is installed typically on top of the reactor. The stripper serves to keep catalyst and excess amounts of the fluorination agent inside the reactor, while allowing product, by-product and HCL to be removed from the reactor as they form.

The reactor and/or stripper may be lined with the fluoropolymer using traditional application techniques such as, for example, spray and bake molding. It has been found, however, that inserting a loose, fluoropolymer liner provides for improved and unexpected results. As used herein, a "loose, fluoropolymer liner" broadly refers to a liner which covers at least portion of metallic part of the reactor and which is fitted from a film or sheet of a fluoropolymer material. Preferably, the sheet has a thickness of no less than about 0.7 mm which is thicker generally than molded liners. As mentioned above, this method of applying the liner is preferred since it tends to produce a less porous lining compared to molding methods like rotary baking. A loose liner is preferred also because it is not limited in thickness and heavy liners may be used. Thicker liners not only lower porosity, but also increase strength. Consequently, a loose, fluoropolymer liner tends to have sufficient strength to span reactor weep holes under reaction pressure. For example, a liner of no less than about 0.7mm has sufficient strength to span at least a 1/4 inch weep hole.

To protect the liner and enhance its reliability, corrosion resistant carbon or silicon carbide brick may be installed in the reactor. These materials are physically more durable than a fluoropolymer liner, and thus are better at resisting abrasion, impact and agitation.

In conducting the reaction, a fluorination agent and a chlorinated organic compound are added to the reactor in the presence of a fluorination catalyst. A halogen exchange then occurs wherein fluorine atoms replace the chlorine atoms of the chlorinated organic compound to produce a desired hydrofluorocarbon.

5

10

As used herein, the term "fluorination agent" refers broadly to any suitable material which provides fluorine for the fluorination reaction. A preferred fluorination agent is substantially anhydrous hydrogen fluoride (HF). Any water that is present in the reaction will tend to react with and deactivate the fluorination catalyst. Therefore, substantially anhydrous HF is preferred. The term "substantially anhydrous" as used herein means that the HF contains less than about 0.1 weight % water and preferably contains less than about 0.05 weight % water. It should be understood, however, that the presence of water in the reaction can be compensated for by increasing the amount of catalyst used.

15

20

25

As used herein, the term "chlorinated organic compound" refers broadly to any hydrocarbon having at least one chlorine atom. Suitable chlorinated organic compounds include hydrochlorofluorocarbons (compounds containing carbon, chlorine and hydrogen), hydrochlorocarbons (compounds containing carbon, chlorine and hydrogen) and chlorofluorocarbons (compounds containing carbon, chlorine and fluorine) or mixtures thereof. In a preferred embodiment, the chlorinated organic compound has a formula of  $C_nH_mF_xCl_y$ , wherein  $1 \le n \le 7$ ,  $0 \le m \le 16$ ,  $1 \le x \le 16$ ,  $0 \le y \le 16$ , and  $(m+x+y) \le (2n+2)$ , more preferably,  $2 \le n \le 5$ ,  $0 \le m \le 12$ ,  $1 \le x \le 12$ ,  $0 \le y \le 12$ , and still more preferably, n = 3,  $0 \le m \le 8$ ,  $1 \le x \le 8$ ,  $0 \le y \le 8$ .

It should be understood that as the number of carbon atoms in the chlorinated organic compound molecule increases, the temperature at which it decomposes generally decreases. Chlorinated organic compounds having two or more carbons tend to decompose at temperatures near or below their boiling point. For such compounds, it is therefore desirable to maintain and react them in their liquid state.

The term "fluorination catalyst", as used herein, means any inorganic metal catalyst used for the substitution of fluorine for chlorine in the chlorinated organic compound. The fluorination catalysts employed are halides of metals such as, for example, Sb (V), Ta (V), Mo(V), Nb (V), Sn (IV), Ti (IV), Sb(III) and B(III) and mixtures of two or more thereof. Preferred catalysts include SbCl<sub>5</sub>, SbF<sub>5</sub>, SbCl<sub>3</sub>, TaCl<sub>5</sub>, SnCl<sub>4</sub>, NbCl<sub>5</sub>, TiCl<sub>4</sub>, MoCl<sub>5</sub>, and mixtures of two or more thereof. Antimony pentachloride is more preferred because of its low cost and availability. The fluorination catalysts used in this invention preferably have a purity of at least about 97%.

10

15

5

The amount of catalyst used can vary widely and can be determined by someone skilled in the art without undue experimentation. The amount depends on a number of factors including the catalyst employed, reactants and other process variables. Although relatively low catalyst concentrations are effective (e.g. less than 0.05 mole per mole organic added), higher quantities (0.1 to 0.5 mole catalyst per mole of organic) may be desirable in order to enhance the reaction rate and consequently improve product output. In a batch process, the more preferred amount of catalyst used is 0.1 to 0.25 mole of catalyst per mole of organic.

20

The chlorinated organic compound and the fluorination agent are added to the reactor which is preferably charged with a fluorination catalyst. It is preferable for the chlorinated organic compound to be fed into the reactor in liquid phase since hydrochlorocarbons with 2 or more carbons tend to decompose at temperatures near their boiling point. Once the reaction is under way, the starting materials may be added under pressure to supply additional reactants for a continuous process.

25

The amount of fluorination agent supplied to the reactor should be at least stoichiometric in relation to fluorine to chlorine, e.g., at least about 6 moles HF per mole of CCl<sub>3</sub> CH<sub>2</sub> CCl<sub>3</sub> or at least about 8 moles HF per mole of CCl<sub>3</sub> CH<sub>2</sub> CCl<sub>2</sub>-CH<sub>2</sub> CCl<sub>3</sub>. Generally, a stoichiometric excess of HF is preferred so that HF essentially acts as a solvent for the fluorination reaction. Excessive amounts of HF, however, limit the output of the product for a batch process. The

HF/organic mole ratio is preferably between about 1:1 and about 30:1; more preferably between about 8:1 and about 20:1; most preferably between about 10:1 and about 15:1.

It may be advantageous to periodically regenerate catalyst due to the deactivation of the catalyst over time. For example, if the catalyst is pentavalent, it is preferable to oxidize the catalyst from its trivalent to its more active pentavalent state. This may be accomplished by any means known in the art. When Sb(V) halides are used as the catalyst, chlorine may be co-fed in an amount sufficient to maintain the antimony salts in the +5 oxidation state which generally equates to about 0.06-0.1 lbs chlorine for every pound of SbCl<sub>5</sub> catalyst. The chlorine may be continuously added when operating in a continuous mode, or periodically added when operating in a batch mode. One of ordinary skill in the art can readily determine without undue experimentation the amount of chlorine to be added in order to optimize the use of the catalyst.

15

20

25

30

5

10

Preferably, the fluorination agent and chlorinated organic compound starting material are simultaneously fed to the reactor after the reactor reaches desired temperature. The temperature at which the fluorination reaction is conducted and the period of reaction will depend on the starting material and catalyst used. One of ordinary skill in the art can readily optimize the conditions of the reaction without undue experimentation to obtain the desired results. Temperatures will generally range from about 30° to about 200°C, preferably from about 50° to about 150°C, and most preferably from about 70 to about 130°C. To add heat to the reaction, it may be beneficial to superheat the fluorination agent and/or to preheat the chlorinated organic compound feeds. Top catalyst stripper temperature should be maintained such that most of the unreacted fluorination agent and catalyst are refluxed to the reactor which depends upon conditions such as reaction temperature, pressure, type of organics, and type of catalyst.

Reaction pressure can vary and optimal pressures can be determined by someone skilled in the art without undue experimentation. Convenient operating

pressure range from about 30 to about 300 psi, preferably from about 60 to about 275 psi, and most preferably from about 70 to about 260 psi.

Reaction times are dependent on several factors including catalyst concentration, the type of catalyst, and the temperature. For a batch process, the progress of the reaction can be monitored conveniently by the increase in pressure due to the formation of by-product HCL. Typical reaction times range from about 1 to about 25 hours, and preferably from about 2 to about 8 hours. For a continuous process, the reaction times ranges from about 1 second to about 5 hours, and, preferably, from about 10 seconds to about 1 hour.

10

15

20

5

In accordance with this invention, HFCs may be produced in high yields and purity with low reactor corrosion by using a fluoropolymer-lined reactor. In the preferred embodiment, HFCs having 2 or more carbon atoms are produced, and in a more preferred embodiment, HFC propanes are produced. Still a more preferred embodiment produces pentafluoropropane and hexafluoropropane, and the most preferred embodiment produces 1,1,1,3,3-pentafluoropropane 1,1,1,3,3,3-hexafluoropropane, and 3,3,3-trifluoropropene.

Examples

The following examples are illustrative of the practice of the present invention.

### Example 1

This example shows a process for producing a hydrofluorocarbon, specifically 1,1,1,3,3 pentafluoropropane (245fa), using a fluoropolymer-lined reactor, specifically a 2.5 gal lined reactor. The reactor was charged with about 19 lb antimony pentachloride catalyst and about 3 lb of liquid anhydrous HF reactant fluorination agent. The reactor was heated subsequently to about 83°C. Next, the organic feed liquid 1,1,1,3,3-pentachloropropane and additional liquid anhydrous HF were fed continuously to the reactor at 0.18 lb/hr and 0.14 lbs/hr

25

• :

respectively. Both reactants were fed at room temperature. The reactor pressure was adjusted to 150 psig.

To recover the HFC, a catalyst stripper was employed. The catalyst stripper consists of a packed column and a condenser installed atop the reactor to keep catalyst and excess amounts of HF inside the reactor, while removing product, by-product, and HCl from the reactor as they form. The top temperature of the catalyst stripper was maintained at about 29°C. The 245fa exiting the catalyst stripper had a purity of about 95%. The productivity of 245fa was about 0.12 lbs/hr. Corrosion was prevented by using PTFE-lined reactor.

10

15

5

#### Example 2

This example shows also a process for producing 245fa using a PTFE-lined reactor, but higher quantities of product and under different operating conditions. A 50 gal PTFE-lined reactor was charged with about 350 lbs antimony pentachloride catalyst. Reactor temperature was raised to 95°C. Liquid 1,1,1,3,3-pentachloropropane (HCC-240) preheated to about 90°C and liquid HF super-heated to 100-150°C were added to the reactor at the rate of 605 lbs/day and 332 lbs/day, respectively. Additionally, 36 lbs/day of chlorine were continuously fed to the reactor to keep catalyst active. The reactor pressure was maintained at about 150 psig.

20

From the product stream, the catalyst recovered HFC-245fa, HF, HCl and organic by-products such as 1,3,3-tetrafluoropropene, 1-chloro-3,3,3-trifluoropropene, and 1-chloro-1,3,3,3-tetrafluoropropene. About 21 lbs/hr pure HCl were recovered from the product steam by low temperature distillation. The yield of 245fa was about 90%. Corrosion of the reactor was prevented by using the PTFE-loose liner.

25

#### Example 3

This example shows a process for producing HFC-236a using a PTFE-lined reactor. In this example, the conditions and equipment used were the same as in

Example 2, except the chlorinated organic compound was changed to 1,1,1,3,3,3-hexachloropropane (HCC-230) and the amount of HF was adjusted to 400 lbs/day. The product steam comprised HCl, 1,1,1,3,3,3-hexafluoropropane (HFC-236fa), 1-chloro-1,1,3,3,3-pentafluoropropane (HCFC-235fa), and 1,1,3,3,3-pentafluoropropene among other components. The latter two components can be recycled to produce more HFC-236a. The overall yield of 236fa was about 90%. Corrosion was again prevented by use of a PTFE-lined reactor.

#### Example 4

10

15

20

25

5

This example illustrates the use of a fluoropolymer loose liner in conducting a reaction in which gaseous reactants are fed to the reaction.

To a 4-gal PTFE-lined reactor, 4.3 lbs antimony pentachloride catalyst and about 15 lbs methylene chloride were charged. The reactor was brought to 110°C with chlorine feed at about 0.1 lb/hr. Then gaseous anhydrous HF and gaseous methylene chloride were fed to the reactor at 0.4 and 0.9 lbs/hr respectively. Both reactants were vaporized before they reached the reactor. The reactor pressure was maintained at 135 psig. The product therein consisted of trifluoromethane (0.19%), difluoromethane (96%), and chlorofluoromethane (3.6%). The productivity of difluoromethane was about 0.5 lbs/hr. Corrosion was prevented by using a PTFE-lined 4-gal reactor.

### Comparative Example

This example compares the corrosion rates of prior art reactor metals to the fluoropolymer-lined reactor of the present invention. Rather than generating corrosion rate data based on actual reactor corrosion, representative coupons of the prior art alloys were used. These coupons were exposed to the same reaction conditions as described in Examples 1-4 above. Coupon thickness was measured before and after each reaction and the difference was calculated. The difference in the coupon thickness was then converted to mils per year (MPY) of corrosion

(herein referred to as the "corrosion rate") based upon 8,000 hr/year operation. The corrosion rates for the coupons subjected to the conditions of Examples 1 and 2 are presented in Tables 1 and 2 respectively. The corrosion rates for the coupons subjected to the conditions of Examples 3 and 4 are substantially similar to those presented in Tables 2 and 1 respectively and are not repeated herein.

TABLE 1--Corrosion Rates Under the Conditions of Example 1

 Material
 Average MPY

 Carbon Steel 1018
 190

 Stainless Steel 316
 130

 Monel 400
 130

 Inconel-600
 210

 20CB3
 260

 Hastelloy C-276
 180

TABLE 2--Corrosion Rates Under the Conditions of Example 2

Average MPY Material 341 Carbon Steel 256 Stainless Steel 316 652 Alloy 20 662 Incoloy 825 701 Inconel 600 190 Hastelloy C-276 368 Monel 400

These tables indicate very high corrosion rates for metals that were exposed to the same conditions as the present invention's fluoropolymer lined reactor which showed little or no corrosion. Generally, a corrosion rate of more than about 10 mil per year is considered to be unacceptable for commercial use. Therefore, none of the alloys tested is acceptable for commercial use under the reaction conditions of the examples.

-12-

10

15

5

20

25

#### **CLAIMS**

### WHAT IS CLAIMED IS

10

15

25

1. A process for producing a hydrofluorocarbon (HFC) comprising the steps of:

providing reactor means for conducting a fluorination reaction wherein at least a portion of said reaction means is lined with a fluoropolymer;

adding a chlorinated organic compound in liquid phase and a fluorination agent to said reactor; and

reacting at least a portion of said chlorinated organic compound with at least a portion of said fluorination agent to produce said HFC.

- 2. The process of claim 1, wherein said fluorination agent is hydrogen fluoride, and said chlorinated organic compound has the formula  $C_nH_mCl_xF_y$ , where  $1 \le n \le 7$ ,  $0 \le m \le 16$ ,  $1 \le x \le 16$ ,  $0 \le y \le 16$ , and  $(m+x+y) \le (2n+2)$ , and wherein a stoichiometric excess of said HF is maintained.
- 3. The process of claim 2, wherein  $3 \le n \le 5$ .
- 4. The process of claim 1, wherein said reactor means comprises a reactor having a loose, fluoropolymer liner.
  - 5. The process of claim 1, wherein said fluoropolymer is selected from the group consisting of polytetrafluoroethylene, poly(vinylidene fluoride), ethylene-tetrafluoroethylene polymer, ethylene-hexafluoropropylene polymer, tetrafluoroethylene-hexafluoropropylene polymer, any modified version thereof, and blends of two or more thereof.

6. The process of claim 1, wherein said fluorinating agent is superheated and said chlorinated organic compound is preheated prior to adding to the reactor.

- 7. A process for producing a hydrofluorocarbon (HFC) comprising the steps of:
  - providing a reactor having at least one weep hole and a fluoropolymer liner; and
  - reacting at least a portion of a chlorinated organic compound with at least a portion of a fluorination agent in said reactor to produce said HFC.

10

15

- 8. The process of claim 7, wherein said fluoropolymer liner is a loose, fluoropolymer liner comprised of a material selected from the group consisting of polytetrafluoroethylene, poly(vinylidene fluoride), ethylene-tetrafluoroethylene polymer, ethylene-hexafluoropropylene polymer, tetrafluoroethylene-hexafluoropropylene polymer, any modified version thereof, and blends of two or more thereof.
- 9. A process for producing a hydrofluorocarbon (HFC) comprising the steps of:
- providing a reactor with a loose, fluoropolymer liner; and reacting at least a portion of a chlorinated organic compound with at least a portion of a fluorination agent in said reactor to produce said HFC.
- 10. The process of claim 9, wherein said fluoropolymer is selected from the group consisting of polytetrafluoroethylene, poly(vinylidene fluoride), ethylene-tetrafluoroethylene polymer, ethylene-hexafluoropropylene polymer,

tetrafluoroethylene-hexafluoropropylene polymer, any modified version thereof, and blends of two or more thereof.

- 11. The process of claim 10, wherein said fluoropolymer is polytetrafluoroethylene or its modified version.
- 12. The process of claim 9, wherein said chlorinated organic compound has the formula  $C_nH_mCl_xF_y$ , where  $1 \le n \le 7$ ,  $0 \le m \le 16$ ,  $1 \le x \le 16$ ,  $0 \le y \le 16$ , and  $(m+x+y) \le (2n+2)$ , and wherein a stoichiometric excess of said HF is maintained in said reactor.
- 13. The process of claim 9, wherein said reactor has a physical size of at least 1000 gallons.
- 14. A process for extending the useful life of a reactor in a method for producing a hydrofluorocarbon (HFC), said method comprising the step of:

performing a fluorination reaction by reacting a chlorinated organic compound and a fluorination agent in the presence of a fluorination catalyst;

said process comprising:

5

10

using a reactor with a loose, fluoropolymer liner to perform said fluorination reaction.

15. The process of claim 14, wherein the corrosion rate of said reactor is less than about 10 mils per year based upon about 8,000 h/yr operation.

16. The process of claim 14, wherein said fluorination reaction further comprises:

- charging said reactor with a liquid phase catalyst selected from the group consisting of halides of Sb, Ta, Sn, Nb, Ti, and Mo, and combinations of two or more thereof;
- feeding said reactor with a chlorinated organic compound in liquid state and hydrogen fluoride (HF), wherein said chlorinated organic compound has the formula  $C_nH_mCl_xF_y$ , where  $1 \le n \le 5$ ,  $0 \le m \le 12$ ,  $1 \le x \le 12$ ,  $0 \le y \le 12$ , and  $(m+x+y) \le (2n+2)$ , and wherein a stoichiometric excess of HF is maintained; and

heating said reactor between about 30°C and about 200°C; pressurizing said reactor between about 30 psig and about 300 psig; and recovering said HFC from said reactor using a catalyst stripper.

- 15 17. The process of Claim 16, wherein said reactor has at least one weep hole.
  - 18. The process of Claim 17, wherein said fluorination reaction further comprises:

adding chlorine to said reactor to regenerate said catalyst.

20

5

- 19. The process of Claim 18, wherein the mole ratio of HF to chlorinated organic compound is about 30:1 to about 1:1.
- 20. The process of claim 19, wherein said fluoropolymer is selected from the group consisting of polytetrafluoroethylene, poly(vinylidene fluoride), ethylene-tetrafluoroethylene polymer, ethylene-hexafluoropropylene polymer,

tetrafluoroethylene-hexafluoropropylene polymer, any modified version thereof, and blends of two or more thereof.

# INTERNATIONAL SEARCH REPORT

Inter. Snal Application No PCT/US 98/12951

A. CLASSII IPC 6	FICATION OF SUBJECT MATTER C07C17/20 C07C19/08 B01J3/	04 B01J19/02	•	
	o International Patent Classification (IPC) or to both national class	ification and IPC	······································	
	SEARCHED  ocumentation searched (classification system followed by classific	cation symbols)		
IPC 6	CO7C B01J			
Documentat	tion searched other than minimum documentation to the extent the	at such documents are included in the fields se	arched	
Electronic d	ata base consulted during the international search (name of data	base and, where practical, search terms used		
C. DOCUMI	ENTS CONSIDERED TO BE RELEVANT			
Category *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.	
Y	CHEMICAL ABSTRACTS, vol. 123, r 18 December 1995 Columbus, Ohio, US; abstract no. 339132, KIMURA M ET AL: "Liquid-phase fluorination of organic compour fluoropolymer reactors" XP002077106 cited in the application see abstract	nds in	1-20	
	& JP 07 233102 A (ASAHI GLASS (LTD; JAPAN) 5 September 1995			
		-/		
X Furt	her documents are listed in the continuation of box C.	Patent family members are listed	in annex.	
• Special ca	ategories of cited documents :	"T" later document published after the inte	rnational filing date	
consid	ent defining the general state of the art which is not dered to be of particular relevance document but published on or after the International date	or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention  "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to		
which citatio	ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another in or other special reason (as specified) tent referring to an oral disclosure, use, exhibition or	involve an inventive step when the do "Y" document of particular relevance; the cannot be considered to involve an indocument is combined with one or m	ocument is taken alone claimed invention eventive step when the ore other such docu-	
"P" docum	means ent published prior to the international filing date but han the priority date claimed	ments, such combination being obvious in the art.  "&" document member of the same patent	·	
Date of the	actual completion of theinternational search	Date of mailing of the international sea	arch report	
1	O September 1998	22/09/1998		
Name and	mailing address of the ISA  European Patent Office, P.B. 5818 Patentlaan 2	Authorized officer		
	NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Bonnevalle, E		

## INTERNATIONAL SEARCH REPORT

Inter. .onal Application No
PCT/US 98/12951

		PCT/US 98/12951
C.(Continua	ition) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	PATENT ABSTRACTS OF JAPAN vol. 015, no. 377 (C-0870), 24 September 1991 & JP 03 153641 A (CENTRAL GLASS CO LTD), 1 July 1991 see abstract	1-20
Y	US 2 875 254 A (F.J. GRADISHAR) 24 February 1959 see column 1 - column 3	1-20
Y	WO 97 05089 A (DU PONT ) 13 February 1997 see page 20, line 29 - line 30; examples 16-20	1-20
Y	GB 656 793 A (COMPAGNIE DE PRODUITS CHIMIQUES ET ELECTRO-METALLURGIQUES) 5 September 1951 see the whole document	1-20
P, X	WO 98 00378 A (ALLIED SIGNAL INC) 8 January 1998 see the whole document	14,16, 18-20
A	US 5 236 013 A (DAS DILIP K) 17 August 1993 see column 2 - column 3	1,4,7
		•
	·	

## INTERNATIONAL SEARCH REPORT

Information on patent family members

Inte. .onal Application No
PCT/US 98/12951 —

Patent document cited in search report		Publication date	Patent family member(s)		Publication date
US 2875254	Α	24-02-1959	FR	1211350 A	15-03-1960
WO 9705089	Α	13-02-1997	AU AU WO	6643696 A 6685796 A 9705090 A	26-02-1997 26-02-1997 13-02-1997
GB 656793	A		NONE		
WO 9800378	Α	08-01-1998	US	5763706 A	09-06-1998
US 5236013	A	17-08-1993	NONE		